08/249689 attribut to Paper # 28

(FILE 'HOME' ENTERED AT 09:35:58 ON 17 AUG 95)

FILE 'CAPLUS' ENTERED AT 09:36:03 ON 17 AUG 95

L1 97 S ANTIBIOTIC AND RNA (2A) BIND?

L2 1 S L1 AND GROOVE

L3 64 S L1 NOT POLYMERASE

=> s 13 range 1985-1990

MISSING OPERATOR 'L3 RANGE'

The search profile that was entered contains terms or nested terms that are not separated by a logical operator.

=> s l3 range=1985-1990

9051 POLYMERASE

L4 6 L1 NOT POLYMERASE

=> d 14 1-6 bib ab

L4 ANSWER 1 OF 6 CAPLUS COPYRIGHT 1995 ACS

AN 1990:212722 CAPLUS

DN 112:212722

TI Translational repression by bacteriophage MS2 coat protein expressed from a plasmid. A system for genetic analysis of a protein-RNA interaction

AU Peabody, David S.

CS Sch. Med., Univ. New Mexico, Albuquerque, NM, 87131, USA

SO J. Biol. Chem. (1990), 265(10), 5684-9 CODEN: JBCHA3; ISSN: 0021-9258

DT Journal

LA English

The coat protein of phage MS2 is a translational repressor. It inhibits the synthesis of the viral replicase by \*\*\*binding\*\*\* a specific \*\*\*RNA\*\*\* structure that contains the replicase translation initiation region. In order to begin a genetic dissection of the repressor activity of coat protein, a 2-plasmid system has been constructed that expresses coat protein and a replicase-.beta.-galactosidase fusion protein from different, compatible plasmids contg. different \*\*\*antibiotic\*\*\* -resistance determinants. The coat protein expressed from the first plasmid (pCT1) represses synthesis of a replicase-.beta.-galactosidase fusion protein encoded on the other plasmid (pRZ5). Mutations in

the translational operator or in coat protein result in constitutive This permits the straightforward isolation synthesis of the enzyme. of mutations in the coat sequence that affect repressor function. Because of the potential importance of cysteine residues for \*\*\*binding\*\*\* , mutations were constructed that substitute serines for the cysteine residues normally present at positions 46 and 101. Both of these mutations result in translational repressor defects. Chromatog. and electron microscopic analyses indicate that the plasmid-encoded wild-type coat protein forms capsids in vivo. The ability of the mutants to adopt and/or maintain the appropriate conformation was assayed by comparing them to the wild-type protein for their ability to form capsids. Both mutants exhibited evidence of improper folding and/or instability, as indicated by their aberrant elution behavior on a column of Sepharose CL-4B. Methods were developed for the rapid purifn. of plasmid-encoded coat protein, facilitating future biochem. analyses of mutant coat proteins.

- L4 ANSWER 2 OF 6 CAPLUS COPYRIGHT 1995 ACS
- AN 1990:31328 CAPLUS
- DN 112:31328
- TI AbrB, a regulator of gene expression in Bacillus, interacts with the transcription initiation regions of a sporulation gene and an \*\*\*antibiotic\*\*\* biosynthesis gene
- AU Robertson, Jeffrey B.; Gocht, Martin; Marahiel, Mohamed A.; Zuber, Peter
- CS Dep. Bot. Microbiol., Oklahoma State Univ., Stillwater, OK, 74078, USA
- SO Proc. Natl. Acad. Sci. U. S. A. (1989), 86(21), 8457-61 CODEN: PNASA6; ISSN: 0027-8424
- DT Journal
- LA English
- AΒ The abrB gene of B. subtilis is believed to encode a repressor that controls the expression of genes involved in starvation-induced processes such as sporulation and the prodn. of antibiotics and Two such genes, spoVG, a sporulation gene of degradative enzymes. B. subtilis, and tycA, which encodes tyrocidine synthetase I of the tyrocidine biosynthetic pathway in B. brevis, are neg. regulated by To exam. the role of abrB in the repression of abrB in B. subtilis. gene transcription, the AbrB protein was purified and then tested for its ability to bind to spoVG and tycA promoter DNA. mobility shift expt., AbrB was found to bind to a DNA fragment contg. the sequence from -95 to +61 of SpoVG. AbrB protein exhibited reduced affinity for DNA of 2 mutant forms of the spoVG promoter that had been shown to be insensitive to abrB-dependent repression in vivo. These studies showed that an upstream A + T-rich sequence from -37 to -95 was required for optimal AbrB

binding. AbrB protein was also obsd. to bind to the tycA gene within a region between the transcription start site and the tycA coding sequence as well as to a region contg. the putative tycA promoter. These findings reinforce the hypothesis that AbrB represses gene expression through its direct interaction with the transcription initiation regions of genes under its control.

- L4 ANSWER 3 OF 6 CAPLUS COPYRIGHT 1995 ACS
- AN 1987:95365 CAPLUS
- DN 106:95365
- TI Involvement of specific portions of ribosomal RNA in defined ribosomal functions: a study utilizing antibiotics
- AU Cundliffe, E.
- CS Dep. Biochem., Univ. Leicester, Leicester, UK
- SO Struct., Funct., Genet. Ribosomes, ["Ribosome Conf."] (1986),
  Meeting Date 1985, 586-604. Editor(s): Hardesty, Boyd; Kramer,
  Gisela. Publisher: Springer, New York, N. Y.
  CODEN: 55HZA6
- DT Conference; General Review
- LA English
- AB A review with 37 refs. on the binding of ribosomes by antibiotics in relation to the role of rRNA in ribosomal function.
- L4 ANSWER 4 OF 6 CAPLUS COPYRIGHT 1995 ACS
- AN 1986:583365 CAPLUS
- DN 105:183365
- TI The binding of the antitumor \*\*\*antibiotic\*\*\* chartreusin to a poly(dA-dT).poly(dA-dT), poly(dG-dC).poly(dG-dC), calf thymus DNA, transfer RNA, and ribosomal RNA
- AU Krueger, William C.; Pschigoda, Loraine M.; Moscowitz, Albert
- CS Upjohn Co., Kalamazoo, MI, 49001, USA
- SO J. Antibiot. (1986), 39(9), 1298-303 CODEN: JANTAJ; ISSN: 0021-8820
- DT Journal
- LA English
- Chartreusin (I) [6377-18-0] binds cooperatively to the poly(dA-dT) [26966-61-0] duplex and the poly(dG-dC) [36786-90-0] duplex. Both the site-exclusion model and the specific site model yield cooperative binding consts. of about 5 .times. 105 M-1 and 3 .times. 105 M-1 for the AT and GC polymers, resp., and the same stoichiometry and intrinsic binding const. for both polymers of 5 nucleosides per binding site and 3.1 .times. 104 M-1. The Scatchard plot for calf thymus DNA is curved in the opposite sense from that of cooperative binding. These binding data did not fit the site-exclusion model with the cooperative binding parameter as a variable nor the specific site, neg.-cooperative binding model. The site-exclusion model with a cooperative binding parameter of unity

yielded a binding const. of about 4 .times. 104 M-1 and a stoichiometry of about 5 nucleotides per binding site. The same model for transfer and rRNA yielded binding consts. of 5 .times. 103 M-1 and 7 .times. 103 M-1 and stoichiometries of about 13 and 6 nucleotides per binding site, resp.

- L4 ANSWER 5 OF 6 CAPLUS COPYRIGHT 1995 ACS
- AN 1985:609100 CAPLUS
- DN 103:209100
- TI Binding of coumermycin A1 to nucleic acids: a spectroscopic approach
- AU Masotti, Lanfranco; Palu, G.; Von Berger, J.; Meloni, G. A.
- CS Fac. Med. Surg., Univ. Parma, Parma, 43100, Italy
- SO Proc. Int. Congr. Chemother., 13th (1983), Volume 6, 113/13-113/16. Editor(s): Spitzy, K. H.; Karrer, K. Publisher: Verlag H. Egermann, Vienna, Austria.
  - CODEN: 53XPA8
- DT Conference
- LA English
- The coumarin- and carbohydrate-contg. \*\*\*antibiotic\*\*\*
  coumermycin A1 (I) interacts with linear and closed circular DNAs as
  well as with rRNAs, as indicated by absorption and fluorescence
  spectroscopy. Fluorescence quenching showed that I is rather deeply
  buried within the interior of DNA (40% quenching). The apparent
  binding consts. for the DNAs and for rRNA were 3.8 and 2.6 .times \*\*
  104M-1, resp. The interaction with DNA is preferential for dA-dT \*\*
  sequences, and the binding is probably intercalative.
- L4 ANSWER 6 OF 6 CAPLUS COPYRIGHT 1995 ACS
- AN 1985:180996 CAPLUS
- DN 102:180996
- TI Binding of small molecules to nucleic acids with tertiary structure

٠.,..

- AU Nechipurenko, Yu. D.
- CS Inst. Mol. Biol., Moscow, USSR
- SO Biofizika (1985), 30(2), 231-2 CODEN: BIOFAI; ISSN: 0006-3029
- DT Journal
- LA Russian
- AB A model was developed which allows a description of the binding of antibiotics and dyes to nucleic acids (DNA or RNA) in which different regions are involved in the formation of a certain tertiary structure. Interactions between different segments of nucleic acid may contribute to the internal overall energy of the macromol. The case when the tertiary structure and the conformational energy of the macromol. are altered upon binding of small mols. is considered. These structural changes affect the shape of the binding isotherm of ligand to the nucleic acid.

Relations are obtained which permit detn. of the dependence of the conformational energy on the degree of binding of ligand to nucleic acid.

## => LOGOFF Y

| COST IN U.S. DOLLARS                       | SINCE FILE | TOTAL            |
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| DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) | SINCE FILE | $\mathtt{TOTAL}$ |
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STN INTERNATIONAL LOGOFF AT 09:42:17 ON 17 AUG 95

L19 ANSWER 2 OF 15 CAPLUS COPYRIGHT 1995 ACS

**DUPLICATE 2** 

AN 1994:235427 CAPLUS

DN 120:235427

TI Peptide antibiotics of the tuberactinomycin family as inhibitors of group I intron RNA splicing

AU Wank, Herbert; Rogers, Jeff; Davies, Julian; Schroeder, Renee

CS Inst. Mikrobiol. Genet., Univ. Wien, Vienna, A-1030, Austria

SO J. Mol. Biol. (1994), 236(4), 1001-10 CODEN: JMOBAK; ISSN: 0022-2836

DT Journal

LA English

AB The tuberactinomycins are a group of cyclic peptide antibiotics, which are potent \*\*\*inhibitors\*\*\* of prokaryotic protein synthesis. The authors report the \*\*\*inhibitory\*\*\* effect of viomycin, di-beta, lysyl-capreomycin IIA and auberactinomycin A on group I intron self-splicing. They compete with the guanosine cofactor for the G-binding site located in the conserved core of the intron. They are 100-fold more active than all other competitive \*\*\*inhibitors\*\*\* described so far (dGTP, arginine or streptomycin), inhibiting splicing at concns. between 10 and 50 .mu.M. Mutation of the G-binding site leads to partial resistance, and the \*\*\*inhibitory\*\*\* effect of these \*\*\*drugs\*\*\* is dependent on Mg2+ concn. This suggests that the tuberactinomycins have more than one contact site with the intron \*\*\*RNA\*\*\* : via the G- \*\*\*binding\*\*\* site and via addnl. contacts with the RNA backbone. Positioning the tuberactinomycins in the three-dimensional model of the td intron core suggests that the charged lysyl side-chain (R1) is in contact with the backbone of the P1 helix. Structure/function analyses with various tuberactinomycin analogs with different activities confirm the involvement of this side-chain in inhibition of group I self-splicing. The demonstration of a new class of splicing \*\*\*inhibitors\*\*\*, the peptide antibiotics, illustrates how antibiotics may interact with catalytic RNA.

L19 ANSWER 9 OF 15 CAPLUS COPYRIGHT 1995 ACS

**DUPLICATE 4** 

AN 1981:77518 CAPLUS

DN 94:77518

TI The translocation inhibitor tuberactinomycin binds to nucleic acids and blocks the in vitro assembly of 50S subunits

- AU Yamada, Takeshi; Teshima, Tadashi; Shiba, Tetsuo; Nierhaus, Knud H.
- CS Res. Inst. Microbial Dis., Osaka Univ., Suita, 565, Japan
- SO Nucleic Acids Res. (1980), 8(23), 5767-77 CODEN: NARHAD; ISSN: 0305-1048

86630

DT Journal

LA English

AB Ribosome binding studies were performed with a 14C-labeled deriv. of viomycin, tuberactinomycin O (TUM O)(I) [33137-73-4]. TUM O bound to 30 S and 50 S subunits. The \*\*\*binding\*\*\* component was the \*\*\*RNA\*\*\*, since ribosomal proteins did not bind the \*\*\*drug\*\*\* . Other RNAs such as tRNA, phage RNA (MS2), and homopolynucleotides also bound the \*\*\*drug\*\*\* . Striking differences in the binding capacities of the various homopolynucleotides were found. Poly(U) [27416-86-0] bound strongly, poly(G) [25191-14-4] and poly(C) [30811-80-4] bound intermediately, and poly(A) [24937-83-5] showed a very low binding. DNA also bound TUM O, although with native DNA the binding was weak. Finally the effects of viomycin [32988-50-4] on the assembly in vitro of the 50 S subunit from Escherichia coli were tested. A very strong inhibition was found: when the reconstitution was performed at 0.5 .times. 10-6M viomycin the particles formed sedimented at about 50 S, but showed a residual activity of <10%. The \*\*\*inhibitory\*\*\* power of viomycin with respect to the in vitro assembly is more pronounced than that found in in vitro systems for protein synthesis.

=> s 124 and rna

152995 RNA

L25 5 L24 AND RNA

=> d 125 1-5 all

L25 ANSWER 1 OF 5 CAPLUS COPYRIGHT 1995 ACS

AN 1993:530904 CAPLUS

DN 119:130904

TI The search for structure-specific nucleic acid-interactive drugs: Effects of compound structure on \*\*\*RNA\*\*\* versus DNA interaction strength

AU Wilson, W. David; Ratmeyer, Lynda; Zhao, Min; Strekowski, Lucjan; Boykin, David

CS Dep. Chem., Georgia State Univ., Atlanta, GA, 30303, USA

SO Biochemistry (1993), 32(15), 4098-104 CODEN: BICHAW; ISSN: 0006-2960

DT Journal

LA English

CC 1-3 (Pharmacology)

OS CJACS-IMAGE; CJACS

AB \*\*\*RNA\*\*\* genomes of a no. of pathogenic \*\*\*RNA\*\*\* viruses, such as HIV-1, have extensive folded conformations with imperfect A-form duplexas that are essential for virus function and could serve as targets for structure-specific antiviral drugs. an initial step in the discovery of such drugs, the interactions with \*\*\*RNA\*\*\* of a wide variety of compds., which are known to to DNA in the \*\*\*minor\*\*\* \*\*\*bind\*\*\* \*\*\*groove\*\*\* classical or by threading intercalation, have been evaluated by thermal melting and viscometric analyses. The corresponding \*\*\*RNA\*\*\* and DNA polymers, poly(A).cntdot.poly(U) and poly(dA).cntdot.poly(dT), were used as test systems for anal. of binding strength and selectivity. \*\*\*RNA\*\*\* Compds. that \*\*\*bind\*\*\* exclusively in the \*\*\*minor\*\*\* \*\*\*qroove\*\*\* at AT sequences of DNA (e.g., netropsin, distamycin, and a zinc porphyrin deriv.) do not have significant interactions with Compds. that \*\*\*bind\*\*\* \*\*\*minor\*\*\* in the in AT sequences of DNA but have other favorable \*\*\*groove\*\*\* interactions in GC sequences of DNA (e.g., Hoechst 33258, DAPI, and other arom. diamidines) can have very strong interactions. A group of classical intercalators and a group of intercalators with unfused arom. ring systems contain compds. that \*\*\*RNA\*\*\* . intercalate and have strong interactions with

this time, no clear pattern of mol. structure that favors
\*\*\*RNA\*\*\* over DNA interactions for intercalators has emerged.
Compds. that bind to DNA by threading intercalation generally bind
to \*\*\*RNA\*\*\* by the same mode, but none of the threading
intercalators tested to date have shown selective interactions with
\*\*\*RNA\*\*\*

- ST antiviral \*\*\*RNA\*\*\* interaction structure
- IT Virucides and Virustats
  - ( \*\*\*RNA\*\*\* -interactive, structure in relation to)
- IT Ribonucleic acids

(antiviral agents binding to, structure in relation to)

- IT Molecular structure-biological activity relationship
  - ( \*\*\*RNA\*\*\* -interacting, of antiviral agents)
- IT 61-73-4, Methylene blue 65-61-2, Acridine orange 92-31-9, Toluidine blue 0 92-62-6, Proflavine 100-33-4, Pentamidine 135-49-9, Acridine yellow G 519-23-3, Ellipticine 1404-15-5, Nogalamycin Benzo[c]cinnoline 3546-21-2, Ethidium 6872-73-7, Coralyne 22291-04-9 23214-92-8, Adriamycin 23491-45-4, Hoechst 33258 34089-71-9 34089-72-0 34089-73-1 39389-47-4, Distamycin 40603-58-5, Zn-P 4 47165-04-8, DAPI 48242-71-3, Ni-P 4 65271-80-9, Mitoxantrone 73819-26-8 78186-34-2, Bisantrene 80498-71-1 80498-74-4 108772-82-3 117269-54-2 124959-47-3 133671-66-6 133671-68-8 149691-35-0, R 11645DA · 148711-61-9 148726-12-9 138172-26-6 ( \*\*\*RNA\*\*\* binding by, structure effect on, antiviral design in relation to)
- L25 ANSWER 2 OF 5 CAPLUS COPYRIGHT 1995 ACS
- AN 1993:74667 CAPLUS
- DN 118:74667
- TI Definition of the binding sites of individual zinc fingers in the transcription factor IIIA-5S \*\*\*RNA\*\*\* gene complex

. 1

- AU Clemens, Karen R.; Liao, Xiubei; Wolf, Veronica; Wright, Peter E.; Gottesfeld, Joel M.
- CS Dep. Mol. Biol., Scripps Res. Inst., La Jolla, CA, 92037, USA
- SO Proc. Natl. Acad. Sci. U. S. A. (1992), 89(22), 10822-6 CODEN: PNASA6; ISSN: 0027-8424
- DT Journal
- LA English
- CC 3-4 (Biochemical Genetics)
- AB A series of polypeptides contg. increasing nos. of zinc fingers of Xenopus transcription factor IIIA has been generated and binding to the 5S \*\*\*RNA\*\*\* gene internal control region has been studied in order to elucidate the mode of interaction of the individual fingers with DNA. By using a combination of DNase I footprinting, methylation interference, and differential binding to mixts. of DNA fragment differing in length by single base pairs, the binding sites

for individual fingers have been defined. These results have led to a model for the interaction of transcription factor IIIA with the internal control region in which fingers 1-3 bind in the major groove of the promoter C block, fingers 7-9 bind in the major groove of the A block, and finger 5 binds in the major groove of the intermediate element. Fingers 4 and 6 each \*\*\*bind\*\*\* across the \*\*\*minor\*\*\* \*\*\*groove\*\*\*, spanning these promoter elements.

- ST transcription factor TFIIIA zinc finger binding; rRNA gene TFIIIA zinc finger site
- IT Xenopus

(5 S rRNA gene of, transcription factor TFIIIA zinc finger domains binding to, sites for)

IT Gene, animal

(for 5S rRNA, of Xenepus, transcription factor TFIIIA zinc finger domain binding sites in)

IT Deoxyribonucleic acid sequences

(of 5S rRNA gene internal control region, o Xenopus, transcription factor TFIIIA binding sites in relation to)

- IT Ribonucleic acids, ribosomal
  - (5 S, gene for, of Xenopus, transcription factor TFIIIA zinc finger domain binding sites in)
- IT Genetic element

(ICR (internal control region), in 5 S rRNA gene of Xenopus, binding sites for transcription factor TFIIIA zinc finger domains in)

IT Ribonucleic acid formation factors

(TFIIIA (transcription factor IIIA), zinc fingers of, of Xenopus, binding sites in 5S rRNA gene for)

IT Genetic element

(promoter, of 5 S rRNA gene of Xenopus, transcription factor TFIIIA zinc finger domains binding to, sites for)

IT Conformation and Conformers

(zinc-finger motif, in Xenopus transcription factor TFIIIA, 5 S rRNA gene binding sites for)

- L25 ANSWER 3 OF 5 CAPLUS COPYRIGHT 1995 ACS
- AN 1991:669973 CAPLUS
- DN 115:269973
- TI Molecular recognition between ligands and nucleic acids: DNA binding characteristics of analogs of Hoechst 33258 designed to exhibit altered base and sequence recognition
- AU Rao, K. Ekambareswara; Lown, J. William
- CS Dep. Chem., Univ. Alberta, Edmonton, AB, T6G 2G2, Can.
- SO Chem. Res. Toxicol. (1991), 4(6), 661-9 CODEN: CRTOEC; ISSN: 0893-228X
- DT Journal

- LA English
- CC 1-3 (Pharmacology)
- OS CJACS
- GI Diagram(s) available in offline prints and/or printed CA Issue.
- The DNA binding characteristics of new analogs of Hoechst 33258 (I), AB contg. pyridine and benzoxazole units and designed for altered base specificity, were evaluated using UV, fluorescence, and CD studies. Like Hoechst 33258 the new analogs also \*\*\*bind\*\*\* \*\*\*minor\*\*\* \*\*\*groove\*\*\* of B-DNA in a nonintercalative The interaction of the compds. with poly(dA-dT) is salt fashion. independent. The studies with poly(dA-dT), ctDNA, and poly(dG-dC) indicated a decrease in the relative binding strength of the new analogs to DNAs compared with the parent mol., Hoechst 33258. Compds. II and III showed acceptance of GC bases adjacent to AT base pairs. None of the compds. studied exhibited affinity for A-DNA, \*\*\*RNA\*\*\* , or Z-DNA. Structure-DNA binding double-stranded relationships of the new analogs compared with their parent mol., Hoechst 33258, are discussed.
- ST Hoechst 33253 analog DNA binding structure; conformation DNA ligand binding
- IT Conformation and Conformers
  - (of DNA, Hoechst 33258 and analogs binding in relation to)
- IT Molecular association
  - (of Hoechst 33258 analogs with DNA, conformation and structure in relation to)

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- IT Molecular structure-biological activity relationship (DNA-binding, of Hoschst 33258 analogs)
- IT 23491-44-3 126824-04-2 126824-05-3 126824-06-4 126824-07-5 126824-08-6 126848-06-4 126898-32-6
  - (binding of, to DNA, conformation and structure in relation to)
- IT 26966-61-0 36786-90-0
  - (double-stranded, Hoechst 33258 and analogs binding to, conformation and structure in relation to)
- L25 ANSWER 4 OF 5 CAPLUS COPYRIGHT 1995 ACS
- AN 1989:208833 CAPLUS
- DN 110:208833
- TI Specific activation of open complex formation at an Escherichia coli promoter by oligo(N-methylpyrrolecarboxamide)s: effects of peptide length and identification of DNA target sites
- AU Martello, Pamela A.; Bruzik, James P.; DeHaseth, Pieter; Youngquist, R. Scott; Dervan, Peter B.
- CS Sch. Med., Case West. Reserve Univ., Cleveland, OH, 44106, USA
- SO Biochemistry (1989), 28(10), 4455-61 CODEN: BICHAW; ISSN: 0006-2960
- DT Journal
- LA English

CC 9-15 (Biochemical Methods)
Section cross-reference(s): 3, 6

OS CJACS

AB It was shown that open complex formation by \*\*\*RNA\*\*\* polymerase at a promoter contq. a block substitution of nonalternating AT sequences in the spacer DNA sepg. the contacted -10 and -35 regions could be accelerated by distamycin. No stimulation was obsd. at a promoter with a substitution of alternating AT base pairs in the same region or at the promoter with wild-type spacer. The effect of distamycin [tris(N-methylpyrrolecarboamide), formally a P3] was compared with that of its extended homologs P4, P5, and P6. stimulatory potential of these synthetic oligopeptides that \*\*\*minor\*\*\* \*\*\*bind\*\*\* in the \*\*\*qroove\*\*\* of DNA ranks in the order P4 > distamycin, P5 > P6. The interaction of these peptides with the 3 promoters was studied by monitoring the positions of the promoter DNA protected from methidiumpropyl-EDTA-Fe(II) cleavage in the presence of different concns. of ligand. Apparently, a higher affinity of oligopeptide for the spacer DNA than for the -10 and/or -35 region is a necessary, but not sufficient, condition for stimulation. Different patterns of protected DNA regions are seen with each of the 3 promoters; with distamycin, P4, and P5, a unique arrangement of protected regions is obsd. for the variant contg. nonalternating AT base pairs in its Thus, differences in the ways the minor-groove binders interact with each of the promoter variants account for the obsd. differential stimulation. Apparently, it is a ligand-induced structural change in the nonalternating AT DNA that is responsible for activation of open complex formation at the promoter contg. this substitution.

ST \*\*\*RNA\*\*\* polymerase complex promoter distamycin; Escherichia; promoter complex formation peptide; methylpyrrolecarboxamide promoter complex activation

IT Escherichia coli

(open complex formation at promoter of, activation of, by distamycin and homologs, peptide length and DNA target site in relation to)

IT Gene and Genetic element, microbial

(promoter, pRM, open complex formation at, activation of, of Escherichia coli by distamycin and homologs, peptide length and DNA target site in relation to)

IT 9014-24-8D, \*\*\*RNA\*\*\* polymerase, complexes with Escherichia coli promoter

(formation of open, activation of, by distamycin and homologs, peptide length and DNA target site in relation to)

IT 120229-11-0 120229-12-1 120229-13-2

(genetic promoter contg., open complex formation at, activation of, by distamycin and homologs, peptide length and DNA target

site in relation to)

IT 90138-97-9 120145-57-5 120145-58-6

(open complex formation and Escherichia coli promoter response to, peptide length and DNA target site in relation to)

IT 636-47-5, Distamycin A

(open complex formation at Escherichia coli promoter response to, peptide length and DNA target site in relation to)

- L25 ANSWER 5 OF 5 CAPLUS COPYRIGHT 1995 ACS
- AN 1971:444807 CAPLUS
- DN 75:44807
- TI Effect of a reporter molecule on chromatin template activity
- AU Farber, John; Baserga, Renato; Gabbay, Edmond J.
- CS Sch. Med., Temple Univ., Philadelphia, Pa., USA
- SO Biochem. Biophys. Res. Commun. (1971), 43(3), 675-81 CODEN: BBRCA9
- DT Journal
- LA English
- CC 2 (General Biochemistry)
- AB A reporter mol., said to \*\*\*bind\*\*\* exclusively to the

  \*\*\*minor\*\*\* \*\*\*groove\*\*\* of DNA, does not interfere with the

  transcription of S3 HeLa cell chromatin by an exogenous Escherichia

  coli \*\*\*RNA\*\*\* polymerase. This is in contrast to the marked

  inhibition of chromatin template activity by actinomycin D. This

  suggests that the chromatin proteins regulating transcription by

  \*\*\*RNA\*\*\* polymerase are located in the major groove of DNA.
- ST chromatin transcription \*\*\*RNA\*\*\* polymerase; actinomycin DNA\*\*
  transcription
- IT Proteins

(of chromatin major groove, in template activity regulation)

IT Chromatin

(template activity of, proteins of major groove in regulation of)

## => LOGOFF Y

| COST IN U.S. DOLLARS                       | SINCE FILE | TOTAL   |
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| FULL ESTIMATED COST                        | 21.49      | 120.47  |
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|  | ENTRY      | SESSION |
| CA SUBSCRIBER PRICE                        | -2.21      | -3.97   |
|  |            |         |

STN INTERNATIONAL LOGOFF AT 20:09:13 ON 15 AUG 95

=> -s rna and minor groove

190453 RNA

55807 MINOR

3400 GROOVE

1053 MINOR GROOVE

(MINOR (W) GROOVE)

L1 90 RNA AND MINOR GROOVE

=> s l1 and bind

41895 BIND

L2 10 L1 AND BIND

=> d l2 1-10 all

L2 ANSWER 1 OF 10 MEDLINE

AN 95281599 MEDLINE

TI A peptide interaction in the major groove of \*\*\*RNA\*\*\* resembles protein interactions in the \*\*\*minor\*\*\* \*\*\*groove\*\*\* of DNA.

AU Chen L; Frankel A D

CS Department of Biochemistry and Biophysics, University of California, San Francisco 94141, USA.

NC AI29135 (NIAID) AI08591 (NIAID)

SO Proc Natl Acad Sci U S A, (1995 May 23) 92 (11) 5077-81. Journal code: PV3. ISSN: 0027-8424.

CY United States

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals; Cancer Journals

EM 9509

A 17-amino acid arginine-rich peptide from the bovine AB immunodeficiency virus Tat protein has been shown to \*\*\*bind\*\*\* with high affinity and specificity to bovine immunodeficiency virus transactivation response element (TAR) \*\*\*RNA\*\*\* , making major groove near a bulge. We show contacts in the \*\*\*RNA\*\*\* that, as in other peptide- \*\*\*RNA\*\*\* complexes, arginine and threonine side chains make important contributions to binding but, unexpectedly, that one isoleucine and three glycine residues also are critical. The isoleucine side chain may intercalate into a \*\*\*RNA\*\*\* . Glycine residues may allow hydrophobic pocket in the deeply within the \*\*\*RNA\*\*\* the peptide to \*\*\*bind\*\*\* groove and may help determine the conformation of the peptide. Similar features have been observed in protein-DNA and drug-DNA

complexes in the DNA \*\*\*minor\*\*\* \*\*\*groove\*\*\* , including hydrophobic interactions and binding deep within the groove, suggesting that the major groove of \*\*\*RNA\*\*\* and \*\*\*minor\*\*\* of DNA may share some common recognition features. \*\*\*aroove\*\*\* Check Tags: Comparative Study; Support, U.S. Gov't, P.H.S. CTAmino Acid Sequence Base Sequence Circular Dichroism DNA, Viral: CH, chemistry \*DNA, Viral: ME, metabolism \*Gene Products, tat: CH, chemistry \*Gene Products, tat: ME, metabolism HIV: ME, metabolism \*Immunodeficiency Virus, Bovine: ME, metabolism Molecular Sequence Data Mutagenesis, Insertional Nucleic Acid Conformation Peptide Fragments: CH, chemistry \*Peptide Fragments: ME, metabolism Protein Conformation Protein Denaturation Recombinant Proteins: CH, chemistry Recombinant Proteins: ME, metabolism \*\*\*\*RNA-Binding Proteins: CH, chemistry\*\*\* \*\*\* RNA-Binding Proteins: ME, metabolism\*\*\* \*\*\*\*RNA, Viral: CH, chemistry\*\*\* \*\*\*\*RNA, Viral: ME, metabolism\*\*\* Thermodynamics \*\*\*136628-24-5 (TAR RNA-binding protein) \*\*\* RN0 (DNA, Viral); 0 (Gene Products, tat); 0 (Peptide Fragments); 0 CN(Recombinant Proteins); 0 ( \*\*\*RNA\*\*\* -Binding Proteins); 0 ( \*\*\*RNA\*\*\* , Viral) ANSWER 2 OF 10 MEDLINE L2MEDLINE AN 95249367 Transferring the purine 2-amino group from guanines to adenines in ΤI DNA changes the sequence-specific binding of antibiotics. AU Bailly C; Waring M J Department of Pharmacology, University of Cambridge, UK. CS Nucleic Acids Res, (1995 Mar 25) 23 (6) 885-92. SO Journal code: O8L. ISSN: 0305-1048. ENGLAND: United Kingdom CYDTJournal; Article; (JOURNAL ARTICLE) LA Priority Journals; Cancer Journals FS EM 9508 The proposition that the 2-amino group of guanine plays a critical AB

role in determining how antibiotics recognise their binding sites in DNA has been tested by relocating it, using tyrT DNA derivative molecules substituted with inosine plus 2,6-diaminopurine (DAP). Irrespective of their mode of interaction with DNA, such GC-specific antibiotics as actinomycin, echinomycin, mithramycin and chromomycin find new binding sites associated with DAP-containing sequences and are excluded from former canonical sites containing I.C base pairs. The converse is found to be the case for a group of normally \*\*\*bind\*\*\* AT-selective ligands which in the \*\*\*minor\*\*\* of the helix, such as netropsin: their preferred \*\*\*groove\*\*\* sites become shifted to IC-rich clusters. Thus the binding sites of all these antibiotics strictly follow the placement of the purine 2-amino group, which accordingly must serve as both a positive and negative effector. The footprinting profile of the 'threading' intercalator nogalamycin is potentiated in DAP plus inosine-substituted DNA but otherwise remains much the same as seen with natural DNA. The interaction of echinomycin with sites containing the TpDAP step in doubly substituted DNA appears much stronger than its interaction with CpG-containing sites in natural DNA.

CTCheck Tags: Support, Non-U.S. Gov't Adenine: CH, chemistry \*Antibiotics, Antineoplastic: ME, metabolism \*Antibiotics, Peptide: ME, metabolism Base Sequence Binding Sites Dinucleoside Phosphates: ME, metabolism DNA: CH, chemistry \*DNA: ME, metabolism Guanine: CH, chemistry Inosine: CH, chemistry Intercalating Agents Ligands Molecular Sequence Data \*\*\* RNA, Transfer, Tyr: GE, genetics\*\*\* \*2-Aminopurine: AA, analogs & derivatives 2-Aminopurine: CH, chemistry 2-Aminopurine: ME, metabolism 1904-98-9 (2,6-diaminopurine); 2382-65-2 (cytidylyl-3'-5'-RN guanosine); 452-06-2 (2-Aminopurine); 58-63-9 (Inosine); 73-24-5 (Adenine); 73-40-5 (Guanine); 9007-49-2 (DNA) 0 (Antibiotics, Antineoplastic); 0 (Antibiotics, Peptide); 0 CN (Dinucleoside Phosphates); 0 (Intercalating Agents); 0 (Ligands); 0 , Transfer, Tyr) \*\*\*RNA\*\*\*

L2 ANSWER 3 OF 10 MEDLINE

AN 94316511 MEDLINE

TI The \*\*\*RNA\*\*\* polymerase I transcription factor UBF is a sequence-tolerant HMG-box protein that can recognize structured nucleic acids.

AU Copenhaver G P; Putnam C D; Denton M L; Pikaard C S

CS Biology Department, Washington University, St Louis, MO 63130.

SO Nucleic Acids Res, (1994 Jul 11) 22 (13) 2651-7. Journal code: O8L. ISSN: 0305-1048.

CY ENGLAND: United Kingdom

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals; Cancer Journals

EM 9410

AB Upstream Binding Factor (UBF) is important for activation of ribosomal \*\*\*RNA\*\*\* transcription and belongs to a family of proteins containing nucleic acid binding domains, termed HMG-boxes, with similarity to High Mobility Group (HMG) chromosomal proteins. Proteins in this family can be sequence-specific or highly sequence-tolerant binding proteins. We show that Xenopus UBF can be classified among the sequence-tolerant class. Methylation interference assays using enhancer DNA probes failed to reveal any critical nucleotides required for UBF binding. Selection by UBF of optimal binding sites among a population of enhancer oligonucleotides with randomized sequences also failed to reveal any \*\*\*groove\*\*\* consensus sequence. The \*\*\*minor\*\*\* drugs chromomycin A3, distamycin A and actinomycin D competed against UBF for enhancer binding, suggesting that UBF, like other HMG-box proteins, probably interacts with the \*\*\*minor\*\*\* . UBF also shares with other HMG box proteins the \*\*\*groove\*\*\* ability to \*\*\*bind\*\*\* synthetic cruciform DNA. However, UBF appears different from other HMG-box proteins in that it can \*\*\*RNA\*\*\* (tRNA) and DNA. The both sequence-tolerant nature of UBF-nucleic acid interactions may accommodate the rapid evolution of ribosomal \*\*\*RNA\*\*\* gene sequences.

CT Check Tags: Animal; Support, Non-U.S. Gov't

Base Sequence

Chromomycin A3: PD, pharmacology Dactinomycin: PD, pharmacology Distamycins: PD, pharmacology

\*DNA: ME, metabolism

\*DNA-Binding Proteins: ME, metabolism

Enhancer Elements (Genetics)

\*High Mobility Group Proteins: ME, metabolism

Methylation

Molecular Sequence Data

Nucleic Acid Conformation

\*\*\*\*RNA Polymerase I: ME, metabolism\*\*\*

\*\*\*\*RNA, Transfer: ME, metabolism\*\*\*
\*Transcription Factors: ME, metabolism
Xenopus laevis

RN 50-76-0 (Dactinomycin); 636-47-5 (distamycin A); 7059-24-7 (Chromomycin A3); 9007-49-2 (DNA); \*\*\*9014-25-9 (RNA, Transfer)\*\*\*

- CN EC 2.7.7.- ( \*\*\*RNA\*\*\* Polymerase I); 0 (transcription factor UBF); 0 (Distamycins); 0 (DNA-Binding Proteins); 0 (High Mobility Group Proteins); 0 (Transcription Factors)
- L2 ANSWER 4 OF 10 MEDLINE
- AN 94271753 MEDLINE
- TI Effects of \*\*\*minor\*\*\* \*\*\*groove\*\*\* binding drugs on the interaction of TATA box binding protein and TFIIA with DNA.
- AU Chiang S Y; Welch J; Rauscher F J 3rd; Beerman T A
- CS Department of Experimental Therapeutics, Roswell Park Cancer Institute, Buffalo, New York 14263.
- NC CA16056 (NCI) CA09072 (NCI) CA52009 (NCI)

+

- SO Biochemistry, (1994 Jun 14) 33 (23) 7033-40. Journal code: AOG. ISSN: 0006-2960.
- CY United States
- DT Journal; Article; (JOURNAL ARTICLE)
- LA English
- FS Priority Journals
- EM 9409
- TBP (TATA box binding protein), a general transcription factor AB required for proper initiation of gene expression by \*\*\*RNA\*\*\* \*\*\*minor\*\*\* \*\*\*groove\*\*\* binding drugs. polymerase II, and (MGBs) both interact with DNA within the \*\*\*minor\*\*\* at AT sites. This study has evaluated MGBs as \*\*\*qroove\*\*\* inhibitors of DNA/TBP complex formation by gel mobility shift assays. Our results demonstrate that reversible MGBs (DAPI, distamycin A, Hoechst 33258, and netropsin) are effective inhibitors of the formation of DNA/TBP complex and that distamycin A is the most potent (0.16 microM inhibits TBP complex formation by 50%). CC-1065, a drug that covalently binds to DNA in the \*\*\*groove\*\*\* , is even more active than distamycin A (0.00085 microM inhibits TBP complex formation by 50%). Significantly more CC-1065 (0.009 microM) is required to break up preformed DNA/TBP complex compared to the drug concentration needed to prevent complex formation. In comparison, the order of drug addition has little influence on the ability of reversible MGBs to disrupt DNA/TBP complex. In the presence of TFIIA, a factor that enhances TBP association with DNA, greater drug concentrations (distamycin A and CC-1065, respectively) are needed to disrupt a preformed complex of

DNA/TBP/TFIIA. In comparison to MGBs, drugs capable of binding to DNA by intercalation are generally weaker at blocking TBP complex formation except for hedamycin, which can intercalate and irreversibly \*\*\*bind\*\*\* to DNA and is as effective as reversible MGBs.

CTCheck Tags: Support, Non-U.S. Gov't; Support, U.S. Gov't, P.H.S. Base Sequence

Distamycins: PD, pharmacology

DNA: CH, chemistry

\*DNA: DE, drug effects

\*DNA: ME, metabolism

\*DNA-Binding Proteins: ME, metabolism

Hoe 33258: PD, pharmacology

Indoles: PD, pharmacology

Intercalating Agents: PD, pharmacology

Leucomycins: PD, pharmacology

Molecular Sequence Data

Netropsin: PD, pharmacology

Protein Binding: DE, drug effects

\*Transcription Factors: ME, metabolism

\*TATA Box

- 1438-30-8 (Netropsin); 23491-45-4 (Hoe 33258); 47165-04-8 (DAPI); RN636-47-5 (distamycin A); 69866-21-3 (CC 1065); 9007-49-2 (DNA)
- 0 (transcription factor TFIIA); 0 (Distamycins); 0 (DNA-Binding CN Proteins); 0 (Indoles); 0 (Intercalating Agents); 0 (Leucomycins); 0 (Transcription Factors); 0 (TATA-box-binding protein)
- L2ANSWER 5 OF 10 MEDLINE
- 94163770 MEDLINE AN
- 3T3 NIH murine fibroblasts and B78 murine melanoma cells expressing TIthe Escherichia coli N3-methyladenine-DNA glycosylase I do not become resistant to alkylating agents.
- Imperatori L; Damia G; Taverna P; Garattini E; Citti L; Boldrini L; ΑU D'Incalci M
- CS Istituto di Ricerche Farmacologiche Mario Negri, Milan, Italy.
- Carcinogenesis, (1994 Mar) 15 (3) 533-7. SO

Journal code: C9T. ISSN: 0143-3334.

- CY ENGLAND: United Kingdom
- Journal; Article; (JOURNAL ARTICLE) DT
- LA English
- FS Priority Journals; Cancer Journals
- 9406 EM
- The role of alkylation of the N3 position of adenine in the AB cytotoxicity of alkylating agents in mammalian cells is still undefined. By co-transfecting NIH3T3 murine fibroblast and murine B78 H1 melanoma cells with pSG5tag and pSV2neo, we obtained clones expressing the mRNA of the bacterial tag gene coding for

N3-methyladenine-DNA glycosylase I (Gly I), which specifically repairs N3-methyladenine. The levels of Gly I were 400 times higher in NIH3T3 pSG5tag (clone 3.9.4) and 12-33 times higher in B78 H1 tag clones (2A4, 2A6, 2C3 and 2D1) than in the respective control cells. The sensitivity to alkylating agents was evaluated in tag-expressing cells in comparison with pSG5, pSV2neo co-transfected control cells. As regards the cytotoxic activity of methylating agents (N-methylnitrosourea, N-methyl-N'-nitro-N-nitrosoquanidine, dimethylsulfate and temozolomide) and other alkylators with different structure and different interactions with DNA such as CC-1065 and FCE-24517 ( \*\*\*minor\*\*\* \*\*\*groove\*\*\* \*\*\*bind\*\*\* to N3 of adenine), 4-[bis(2chloroethyl) amino] - L-phenylalanine and cis-diamminedichloroplatinum II, cytotoxicity was the same for tag-expressing and non-expressing cells. These results suggest that the increased expression of N3-methyladenine-DNA glycosylase is not necessarily a crucial mechanism for the resistance of cells to alkylating agents. Check Tags: Animal; Support, Non-U.S. Gov't \*Adenine: AA, analogs & derivatives Alkylating Agents: PD, pharmacology Drug Resistance: GE, genetics \*Escherichia coli: EN, enzymology Escherichia coli: GE, genetics Gene Expression Regulation, Enzymologic \*Genes, Bacterial Melanoma: DT, drug therapy \*Melanoma: EN, enzymology Melanoma: GE, genetics Mice Nucleosidases: GE, genetics \*Nucleosidases: ME, metabolism \*\*\* RNA, Messenger: ME, metabolism\*\*\* Transfection Tumor Cells, Cultured 3T3 Cells: DE, drug effects \*3T3 Cells: EN, enzymology 73-24-5 (Adenine) EC 3.2.2. (Nucleosidases); 0 (Alkylating Agents); 0 ( \*\*\*RNA\*\*\* Messenger) tag ANSWER 6 OF 10 MEDLINE 93229513 MEDLINE The search for structure-specific nucleic acid-interactive drugs: effects of compound structure on \*\*\*RNA\*\*\* versus DNA interaction strength. Wilson W D; Ratmeyer L; Zhao M; Strekowski L; Boykin D

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CS Department of Chemistry, Georgia State University, Atlanta 30303.

NC - AI-27196 (NIAID)

SO Biochemistry, (1993 Apr 20) 32 (15) 4098-104.

Journal code: AOG. ISSN: 0006-2960.

CY United States

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 9307

AB The \*\*\*RNA\*\*\* genomes of a number of pathogenic viruses, such as HIV-1, have extensive folded conformations with imperfect A-form duplexes that are essential for virus function and could serve as targets for structure-specific antiviral drugs. As an initial step in the discovery of such drugs, the interactions with of a wide variety of compounds, which are known to to DNA in the \*\*\*minor\*\*\* \*\*\*groove\*\*\* \*\*\*bind\*\*\* classical or by threading intercalation, have been evaluated by thermal melting and viscometric analyses. The corresponding sequence and DNA polymers, poly(A).poly(U) and poly(dA).poly(dT), were used as test systems for analysis of \*\*\*RNA\*\*\* binding strength and selectivity. Compounds that \*\*\*bind\*\*\* exclusively \*\*\*minor\*\*\* \*\*\*qroove\*\*\* in AT sequences of DNA (e.g., netropsin, distamycin, and a zinc porphyrin derivative) do \*\*\*RNA\*\*\* not have significant interactions with . Compounds that in the minor grove in AT sequences of DNA but have other favorable interactions in GC sequences of DNA (e.g., Hoechsts 33258, DAPI, and other aromatic diamidines) can have very strong interactions. A group of classical intercalators and a group of intercalators with unfused aromatic ring systems contain compounds that intercalate and have strong interactions with . At this time, no clear pattern of molecular structure that favors \*\*\*RNA\*\*\* over DNA interactions for intercalators has emerged. Compounds that \*\*\*bind\*\*\* to DNA by threading \*\*\*RNA\*\*\* intercalation generally \*\*\*bind\*\*\* by the same to mode, but none of the threading intercalators tested to date have shown selective interactions with \*\*\*RNA\*\*\*

Check Tags: Comparative Study; Support, U.S. Gov't, P.H.S.

\*DNA: CH, chemistry

Genome, Viral

CT

HIV-1: GE, genetics

\*Intercalating Agents

Molecular Structure

Nucleic Acid Conformation

\*Poly dA-dT: CH, chemistry

\*Poly A-U: CH, chemistry

\*\*\*\*RNA: CH, chemistry\*\*\*

\*\*\* RNA, Viral: GE, genetics\*\*\*

Structure-Activity Relationship Viscosity

RN 24936-38-7 (Poly A-U); 26966-61-0 (Poly dA-dT); 9007-49-2 (DNA)

- L2 ANSWER 7 OF 10 MEDLINE
- AN 93066335 MEDLINE
- TI Definition of the binding sites of individual zinc fingers in the transcription factor IIIA-5S \*\*\*RNA\*\*\* gene complex.
- AU Clemens K R; Liao X; Wolf V; Wright P E; Gottesfeld J M
- CS Department of Molecular Biology, Scripps Research Institute, La Jolla, CA 92037.
- NC GM36643 (NIGMS) GM26453 (NIGMS) F32 CA09023 (NCI)
- SO Proc Natl Acad Sci U S A, (1992 Nov 15) 89 (22) 10822-6. Journal code: PV3. ISSN: 0027-8424.
- CY United States
- DT Journal; Article; (JOURNAL ARTICLE)
- LA English
- FS Priority Journals; Cancer Journals
- EM 9302
- A series of polypeptides containing increasing numbers of zinc AB fingers of Xenopus transcription factor IIIA has been generated and gene internal control region has \*\*\*RNA\*\*\* binding to the 5S been studied in order to elucidate the mode of interaction of the individual fingers with DNA. By using a combination of DNase I footprinting, methylation interference, and differential binding to mixtures of DNA fragments differing in length by single base pairs, the binding sites for individual fingers have been defined. These results have led to a model for the interaction of transcription factor IIIA with the internal control region in which fingers 1-3 in the major groove of the promoter C block, fingers \*\*\*bind\*\*\* in the major groove of the A block, and finger 5 \*\*\*bind\*\*\* binds in the major groove of the intermediate element. Fingers 4 and \*\*\*bind\*\*\* across the \*\*\*minor\*\*\* \*\*\*groove\*\*\* , spanning these promoter elements.
- CT Check Tags: Animal; Support, Non-U.S. Gov't; Support, U.S. Gov't, P.H.S.

Amino Acid Sequence

Base Sequence

Binding Sites

Cloning, Molecular

\*DNA, Ribosomal: GE, genetics

\*DNA, Ribosomal: ME, metabolism

Escherichia coli: GE, genetics

Methylation Models, Structural Molecular Sequence Data Nucleic Acid Conformation Oligodeoxyribonucleotides Polymerase Chain Reaction: MT, methods Protein Conformation Restriction Mapping \*\*\*\*RNA, Ribosomal, 5S: GE, genetics\*\*\* Transcription Factors: GE, genetics \*Transcription Factors: ME, metabolism Xenopus Zinc Fingers: GE, genetics \*Zinc Fingers: PH, physiology 0 (transcription factor TFIIIA); 0 (DNA, Ribosomal); 0 (Oligodeoxyribonucleotides); 0 ( \*\*\*RNA\*\*\* , Ribosomal, 5S); 0 (Transcription Factors) ANSWER 8 OF 10 MEDLINE 92223348 MEDLINE Molecular recognition between ligands and nucleic acids: DNA binding characteristics of analogues of Hoechst 33258 designed to exhibit altered base and sequence recognition. Rao K E; Lown J W Department of Chemistry, University of Alberta, Edmonton, Canada. Chem Res Toxicol, (1991 Nov-Dec) 4 (6) 661-9. Journal code: A5X. ISSN: 0893-228X. United States Journal; Article; (JOURNAL ARTICLE) English Priority Journals 9207 The DNA binding characteristics of new analogues (2-8) of Hoechst 33258 (1), containing pyridine and benzoxazole units and designed for altered base specificity, were evaluated using UV, fluorescence, and circular dichroism studies. Like Hoechst 33258 the new analogues also \*\*\*bind\*\*\* through the \*\*\*minor\*\*\* \*\*\*qroove\*\*\* B-DNA in a nonintercalative fashion. The interaction of the compounds with poly(dA-dT) is salt independent. The studies with poly(dA-dT), ct DNA, and poly(dG-dC) indicated a decrease in the relative binding strength of the new analogues to DNAs compared with the parent molecule, Hoechst 33258. Compounds 5 and 7 showed acceptance of GC bases adjacent to AT base pairs. None of the compounds studied exhibited affinity for A-DNA, double-stranded , or Z-DNA. Structure-DNA binding relationships of the

new analogues compared with their parent molecule, Hoechst 33258,

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are discussed.

Check Tags: Support, Non-U.S. Gov't Base Sequence Circular Dichroism \*DNA: ME, metabolism \*Hoe 33258: ME, metabolism Nucleic Acid Conformation Osmolar Concentration Poly dA-dT: ME, metabolism Polydeoxyribonucleotides: ME, metabolism Structure-Activity Relationship 23491-45-4 (Hoe 33258); 26966-61-0 (Poly dA-dT); 29855-95-6 RN(poly(dC-dG)); 9007-49-2 (DNA) 0 (Polydeoxyribonucleotides) CNL2ANSWER 9 OF 10 MEDLINE AN 92073376 MEDLINE TIStructural polymorphism in the major groove of a 5S gene complements the zinc finger domains of transcription factor Huber P W; Morii T; Mei H Y; Barton J K ΑU Department of Chemistry and Biochemistry, University of Notre Dame, CS IN 46556. NC GM33309 (NIGMS) CA33620 (NCI) GM38200 (NIGMS) Proc Natl Acad Sci U S A, (1991 Dec 1) 88 (23) 10801-5. SO Journal code: PV3. ISSN: 0027-8424. United States CY DT Journal; Article; (JOURNAL ARTICLE) English LA FS Priority Journals; Cancer Journals EM9203 \*\*\*bind\*\*\* to DNA on the basis of AB Metal complexes that shape-selection have been used to map the conformational features of the DNA binding site for transcription factor IIIA. Conformationally distinct segments are detected on the 5S rRNA gene that correspond closely to the binding sites identified for the individual zinc finger domains of the protein. The local conformations are characterized by a major groove opened because of a change in base pair inclination and/or displacement at a central 5'-pyrimidine-purine-3' step, flanked by a widened \*\*\*groove\*\*\* , as would arise at the junctions between alternating B- and A-like DNA segments. Docking experiments with a consensus structure of a zinc finger reveal that the mixed A-B binding site accommodates the peptide domain better than either canonical B- or

A-DNA helices. The close structural matching of the conformational variations in the 5S rDNA both to the proposed sites of zinc finger

binding and to the shape of an individual zinc finger domain points to DNA structural polymorphism as providing an important determinant in recognition. In particular, shape selection in the 5' half of the internal control region may orient the multiple finger domains. Check Tags: Animal; Support, Non-U.S. Gov't; Support, U.S. Gov't, P.H.S.

Base Sequence
Binding Sites
Computer Simulation
\*Genes, Structural
Models, Molecular
Molecular Sequence Data
Nucleic Acid Conformation
Plasmids
\*Polymorphism (Genetics)

Protein Conformation
Restriction Mapping
\*\*\*\*RNA, Ribosomal, 5S: GE,

\*\*\*\*RNA, Ribosomal, 5S: GE, genetics\*\*\*
\*Transcription Factors: ME, metabolism
Xenopus

\*Zinc Fingers: GE, genetics

- CN 0 (transcription factor TFIIIA); 0 (Plasmids); 0 ( \*\*\*RNA\*\*\* Ribosomal, 5S); 0 (Transcription Factors)
- L2 ANSWER 10 OF 10 MEDLINE
- AN 90344837 MEDLINE
- Detection of drug binding to DNA by hydroxyl radical footprinting, Relationship of distamycin binding sites to DNA structure and positioned nucleosomes on 5S \*\*\*RNA\*\*\* genes of Xenopus.
- AU Churchill M E; Hayes J J; Tullius T D
- CS Department of Chemistry, Johns Hopkins University, Baltimore, Maryland 21218.
- NC CA 37444 (NCI)
  GM 40894 (NIGMS)
  CA 01208 (NCI)
- SO Biochemistry, (1990 Jun 26) 29 (25) 6043-50. Journal code: AOG. ISSN: 0006-2960.
- CY United States
- DT Journal; Article; (JOURNAL ARTICLE)
- LA English
- FS Priority Journals
- EM 9011

CT

AB We report the use of hydroxyl radical footprinting to analyze the interaction of distamycin and actinomycin with the 5S ribosomal \*\*\*RNA\*\*\* genes of Xenopus. There is a qualitative difference in the hydroxyl radical footprints of the two drugs. Distamycin gives a conventional (albeit high-resolution) footprint, while actinomycin

does not protect DNA from hydroxyl radical attack, but instead induces discrete sites of hyperreactivity. We find concentration-dependent changes in the locations of distamycin binding sites on the somatic 5S gene of Xenopus borealis. A high-affinity site, containing a G.C base pair, is replaced at higher levels of bound drug by a periodic array of different lower affinity sites that coincide with the places where the \*\*\*groove\*\*\* of the DNA would face in toward a nucleosome core that is known to \*\*\*bind\*\*\* to the same sequence. These results suggest that distamycin recognizes potential binding sites more by the shape of the DNA than by the specific sequence that is contained in the site and that structures of many sequences are deformable to a shape that allows drug binding. We discuss the utility of hydroxyl radical footprinting of distamycin for investigating the underlying structure of DNA.

CTCheck Tags: Animal; Comparative Study; Support, Non-U.S. Gov't; Support, U.S. Gov't, P.H.S. Actinomycin: ME, metabolism Base Composition Base Sequence Binding Sites Cytosine: ME, metabolism Distamycins: ME, metabolism \*DNA: ME, metabolism Edetic Acid Guanine: ME, metabolism Hydroxides Kinetics Methods Molecular Sequence Data Nucleic Acid Conformation Nucleosomes: PH, physiology \*\*\*\*RNA, Ribosomal: GE, genetics\*\*\* \*\*\*\*RNA, Ribosomal, 5S: GE, genetics\*\*\* \*Xenopus: GE, genetics 1402-38-6 (Actinomycin); 3352-57-6 (Hydroxyl Radical); 60-00-4 RN(Edetic Acid); 71-30-7 (Cytosine); 73-40-5 (Guanine); 9007-49-2 0 (methidiumpropyl-EDTA-iron(II)); 0 (Distamycins); 0 (Hydroxides); CN

=> LOGOFF Y

Ribosomal, 5S)

COST IN U.S. DOLLARS
SINCE FILE TOTAL
ENTRY SESSION
FULL ESTIMATED COST
2.19
2.34

0 (Nucleosomes); 0 ( \*\*\*RNA\*\*\* , Ribosomal); 0 ( \*\*\*RNA\*\*\*

(FILE 'HOME' ENTERED AT 16:29:37 ON 04 AUG 95) 17:17:29 COPY AND CLEAR PAGE, PLEASE

FILE 'MEDLINE, EMBASE, CAPLUS, BIOTECHDS' ENTERED AT 16:34:41 ON 04
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FILE 'MEDLINE'
O SEARCH RNA AND DESIGNER AND INHIBITOR AND REVIEW

L1 Ø SEARCH RNA AND DESIGNER AND INHIBITOR AND REVIEW FILE 'EMBASE'

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L6 Ø SEARCH RNA AND INHIBITOR AND DESIGNER FILE 'EMBASE'

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FILE 'MEDLINE'

L11 78 SEARCH RNA AND RATIONAL FILE 'EMBASE'

L12 88 SEARCH RNA AND RATIONAL FILE 'CAPLUS'

L13 63 SEARCH RNA AND RATIONAL FILE 'BIOTECHDS'

L14 18 SEARCH RNA AND RATIONAL

TOTAL FOR ALL FILES
L15 247 SEARCH RNA AND RATIONAL
FILE 'MEDLINE'

L16 5 SEARCH RNA AND RATIONAL AND REVIEW

FILE 'EMBASE'
L17 31 SEARCH RNA AND RATIONAL AND REVIEW
FILE 'CAPLUS'

L18 12 SEARCH RNA AND RATIONAL AND REVIEW FILE 'BIOTECHDS'

L19 3 SEARCH RNA AND RATIONAL AND REVIEW TOTAL FOR ALL FILES

L20 51 SEARCH RNA AND RATIONAL AND REVIEW SET PAGELENGTH 25

FILE 'MEDLINE'

L21 0 S RNA AND RATIONAL AND INHIBITOR AND REVIEW 17:17:44 COPY AND CLEAR PAGE, PLEASE

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Ø S RNA AND RATIONAL AND INHIBITOR AND REVIEW L24 TOTAL FOR ALL FILES 5 S RNA AND RATIONAL AND INHIBITOR AND REVIEW L25 FILE 'MEDLINE' 78 SEARCH RNA AND RATIONAL L26 FILE 'EMBASE' 88 SEARCH RNA AND RATIONAL L27 FILE 'CAPLUS' 63 SEARCH RNA AND RATIONAL L28 FILE 'BIOTECHDS' L29 18 SEARCH RNA AND RATIONAL TOTAL FOR ALL FILES 247 SEARCH RNA AND RATIONAL L30 FILE 'MEDLINE' 19 SEARCH RNA AND RATIONAL RAN=(1985-1990) L31 FILE 'EMBASE' L32 15 SEARCH RNA AND RATIONAL RAN=(1985-1990) FILE 'CAPLUS' 17:17:53 COPY AND CLEAR PAGE, PLEASE 11 SEARCH RNA AND RATIONAL RAN=(1985-1990) L33 FILE 'BIOTECHDS' L34 6 SEARCH RNA AND RATIONAL RAN=(1985-1990) TOTAL FOR ALL FILES 51 SEARCH RNA AND RATIONAL L35 FILE 'MEDLINE' L36 Ø SEARCH RNA AND RATIONAL AND REVIEW RAN=(1985-1990) FILE 'EMBASE' 5 SEARCH RNA AND RATIONAL AND REVIEW RAN=(1985-1990) L37 FILE 'CAPLUS' L38 3 SEARCH RNA AND RATIONAL AND REVIEW RAN=(1985-1990) FILE 'BIOTECHDS' RAN=(1985-1990) L39 1 SEARCH RNA AND RATIONAL AND REVIEW TOTAL FOR ALL FILES L40 9 SEARCH RNA AND RATIONAL AND REVIEW FILE 'MEDLINE' L41 Ø SEARCH RATIONAL DRUG DESIGN AND RNA RAN=(1985-1990) FILE 'EMBASE' RAN=(1985-1990) L42 O SEARCH RATIONAL DRUG DESIGN AND RNA FILE 'CAPLUS' Ø SEARCH RATIONAL DRUG DESIGN AND RNA RAN=(1985-1990) L43 FILE 'BIOTECHDS' L44 Ø SEARCH RATIONAL DRUG DESIGN AND RNA RAN=(1985-1990) 17:17:58 COPY AND CLEAR PAGE, PLEASE TOTAL FOR ALL FILES L45 O SEARCH RATIONAL DRUG DESIGN AND RNA FILE 'MEDLINE' 5944 S RNA AND INHIBITOR RAN=(ALL) L46 FILE 'EMBASE' L47 4940 S RNA AND INHIBITOR RAN=(ALL) FILE 'CAPLUS' L48 1232 S RNA AND INHIBITOR RAN=(1985-1990) FILE 'BIOTECHDS' L49 267 S RNA AND INHIBITOR RAN=(ALL) TOTAL FOR ALL FILES L50 12383 S RNA AND INHIBITOR

FILE 'MEDLINE'

L51 15 S RNA AND INHIBITOR AND REVIEW

FILE 'EMBASE'

L52 157 S RNA AND INHIBITOR AND REVIEW

FILE 'CAPLUS'

L53 101 S RNA AND INHIBITOR AND REVIEW

FILE 'BIOTECHDS'

L54 6 S RNA AND INHIBITOR AND REVIEW

TOTAL FOR ALL FILES

L55 279 S RNA AND INHIBITOR AND REVIEW

FILE 'MEDLINE'

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L56 5 S RNA AND INHIBITOR AND REVIEW RAN=(1985-1990)

FILE 'EMBASE'

L57 12 S RNA AND INHIBITOR AND REVIEW RAN=(1985-1990)

FILE 'CAPLUS'

L58 18 S RNA AND INHIBITOR AND REVIEW RAN=(1985-1990)

FILE 'BIOTECHDS'

L59 2 S RNA AND INHIBITOR AND REVIEW RAN=(19885-1990)

TOTAL FOR ALL FILES

L60 37 S RNA AND INHIBITOR AND REVIEW

=> d 160 37 abs

L60 ANSWER 37 OF 37 BIOTECHDS COPYRIGHT 1995 DERWENT INFORMATION LTD AN 83-02111 BIOTECHDS 17:18:23 COPY AND CLEAR PAGE, PLEASE

ANSWER 37 OF 37 BIOTECHDS COPYRIGHT 1995 DERWENT INFORMATION LTD L60 An EMBO workshop was held on the replication of prokaryotic DNA. AB Technically there has been a very rapid progress in the understanding of replication control during the last couple of This is due to the appearance of a whole series of new techniques: analysis by restriction endonucleases, cloning of DNA fragments on vectors, DNA and \*\*\*RNA\*\*\* nucleotide sequence analysis, computer-based interpretation of nucleotide sequences, in vitro replication systems, vectors that can be used to analyze for promoters and expression of open reading frames in the nucleotide sequence, etc. Several patterns emerge for replication control by analysis of the basic replicon of several plasmids for DNA nucleotide sequence, promoters, putative genes, transcripts, polypeptides, control functions, etc.: replication control involves at least one plasmid-coded inhibitory function; the target for the is not the origin itself; the inhibitors may be \*\*\*inhibitor\*\*\* small, basic proteins (80-100 aminoacids) or small, unstable molecules (80-110 nucleotides). The workshop gave a useful updating of the current knowledge about replication control. (46 ref)

FILE 'USPAT' ENTERED AT 11:57:28 ON 06 AUG 95

\* WELCOME TO THE

\* U.S. PATENT TEXT FILE

=> s RNA and inhibitor (3A) function

6565 RNA

33242 INHIBITOR

655640 FUNCTION

549 INHIBITOR (3A) FUNCTION

L1 29 RNA AND INHIBITOR (3A) FUNCTION

= > d 11 1-29 bib

US PAT NO: 5,436,321 [IMAGE AVAILABLE] L1: 1 of 29

DATE ISSUED: Jul. 25, 1995

TITLE: Antibodies to the lipopolysaccharide bonding opsonin

septin

INVENTOR: Samuel D. Wright, Larchmont, NY

ASSIGNEE: The Rockefeller University, New York, NY (U.S. corp.)

APPL-NO: 07/916,160 DATE FILED: Jul. 31, 1992

ART-UNIT: 186

PRIM-EXMR: David L. Lacey ASST-EXMR: Susan Loring

LEGAL-REP: Klauber & Jackson

US PAT NO: 5,424,200 [IMAGE AVAILABLE] L1: 2 of 29

DATE ISSUED: Jun. 13, 1995

TITLE: Method for enhanced expression of a DNA sequence of

interest

INVENTOR: Joan C. McPherson, Vancouver, Canada

Robert Kay, Vancouver, Canada

ASSIGNEE: Monsanto Company, St. Louis, MO (U.S. corp.)

APPL-NO: 08/272,900 DATE FILED: Jul. 11, 1994

ART-UNIT: 184

PRIM-EXMR: Patricia R. Moody

LEGAL-REP: Grace L. Bonner, Dennis R. Hoerner, Jr., Richard H. Shear

US PAT NO:

5,424,191 [IMAGE AVAILABLE]

L1: 3 of 29

DATE ISSUED: Jun. 13, 1995

TITLE:

Epithelial cell specific differentiation marker

**INVENTOR:** 

Gaddamanugu L. Prasad, Rockville, MD

Herbert L. Cooper, Rockville, MD

**ASSIGNEE:** 

The United States of America as represented by the

Department of Health and Human Services, Washington, DC

(U.S. govt.)

APPL-NO:

07/887,072

DATE FILED:

May 20, 1992

ART-UNIT:

187

PRIM-EXMR:

Margaret Parr

**ASST-EXMR:** 

Kenneth R. Horlick

LEGAL-REP:

Knobbe, Martens Olson & Bear

US PAT NO:

5,422,344 [IMAGE AVAILABLE]

L1: 4 of 29

DATE ISSUED: Jun. 6, 1995

TITLE:

Method of treating retroviral infections in mammals

**INVENTOR:** 

Esther Priel, Beer Sheva, Israel Donald G. Blair, Kensington, MD

Stephen D. Showalter, Frederick, MD

ASSIGNEE:

The United States of America as represented by the Secretary of the Department of Health & Human Services,

Washington, DC (U.S. govt.)

APPL-NO:

07/520,456

DATE FILED:

May 8, 1990

**ART-UNIT:** 

125

PRIM-EXMR:

Raymond Henley, III

**ASST-EXMR:** 

Russell Travers

LEGAL-REP:

Birch, Stewart, Kolasch & Birch

US PAT NO:

5,403,952 [IMAGE AVAILABLE]

L1: 5 of 29

DATE ISSUED: Apr. 4, 1995

TITLE:

Substituted cyclic derivatives as novel antidegenerative

agents

**INVENTOR:** 

William Hagmann, Westfield, NJ

Charles G. Caldwell, Scotch Plains, NJ

Paul R. Gooley, Westfield, NJ

**ASSIGNEE:** 

Merck & Co., Inc., Rahway, NJ (U.S. corp.)

APPL-NO:

08/133,493

DATE FILED:

Oct. 8, 1993

ART-UNIT:

125

PRIM-EXMR:

Marianne M. Cintins

**ASST-EXMR:** 

Keith MacMillan

LEGAL-REP: Curtis C. Panzer, David L. Rose, Robert J. North

US PAT NO: 5,380,660 [IMAGE AVAILABLE] L1: 6 of 29

DATE ISSUED: Jan. 10, 1995

TITLE: Method of treating serum or serum-containing medium to

inactivate an inhibitor of hepatocyte differentiation

INVENTOR: Douglas M. Jefferson, Watertown, MA

David E. Johnston, Natick, MA

ASSIGNEE: New England Medical Center Hospitals, Inc., Boston, MA

(U.S. corp.)

APPL-NO: 07/956,595

DATE FILED: Oct. 5, 1992

ART-UNIT: 188

PRIM-EXMR: Douglas W. Robinson

ASST-EXMR: Susan M. Dadio LEGAL-REP: Fish & Richardson

US PAT NO: 5,370,991 [IMAGE AVAILABLE] L1: 7 of 29

DATE ISSUED: Dec. 6, 1994

TITLE: Cloned gene encoding human monocyte elastase inhibitor

INVENTOR: Eileen Remold-O'Donnell, Brookline, MA

ASSIGNEE: The Center for Blood Research, Inc., Boston, MA (U.S.

corp.)

APPL-NO: 07/755,461 DATE FILED: Sep. 6, 1991

ART-UNIT: 187

PRIM-EXMR: Amelia Burgess Yarbrough

LEGAL-REP: Wolf, Greenfield & Sacks

US PAT NO: 5,369,125 [IMAGE AVAILABLE] L1: 8 of 29

DATE ISSUED: Nov. 29, 1994

TITLE: Cholesterol-lowering agents

INVENTOR: Gregory D. Berger, Belle Mead, NJ

James D. Bergstrom, Neshanic, NJ

Tesfaye Biftu, Westfield, NJ

Robert L. Bugianesi, Colonia, NJ

Robert M. Burk, Laguna Beach, CA

Narindar N. Girotra, Old Bridge, NJ C. H. Kuo, South Plainfield, NJ

William H. Parsons, Edison, NJ

Mitree M. Ponpipom, Branchburg, NJ

Lori L. Whiting, West Carrollton, OH

ASSIGNEE: Merck & Co., Inc., Rahway, NJ (U.S. corp.)

APPL-NO:

08/033,913

DATE FILED:

Mar. 19, 1993

**ART-UNIT:** 

126

PRIM-EXMR:

Nicky Chan

LEGAL-REP:

Catherine A. Dolan, Melvin Winokur, Paul D. Matukaitis

US PAT NO:

5,364,948 [IMAGE AVAILABLE]

L1: 9 of 29

DATE ISSUED: Nov. 15, 1994

TITLE:

Biologically active compounds isolated from aerobic

fermentation of Trichoderma viride

Guy H. Harris, Cranford, NJ

Deborah Zink, Manalapan, NJ

E. Tracy T. Jones, Solana Beach, CA

Yu L. Kong, Edison, NJ

**ASSIGNEE:** 

**INVENTOR:** 

Merck & Co., Inc., Rahway, NJ (U.S. corp.)

APPL-NO:

08/015,498

DATE FILED:

Feb. 9, 1993

ART-UNIT:

124

PRIM-EXMR:

Jose G. Dees

ASST-EXMR:

Deborah D. Carr

LEGAL-REP:

Catherine A. Dolan, Melvin Winokur, Paul D. Matukaitis

US PAT NO:

5,359,142 [IMAGE AVAILABLE]

L1: 10 of 29

DATE ISSUED: Oct. 25, 1994

TITLE:

Method for enhanced expression of a protein

**INVENTOR:** 

Joan C. McPherson, Vancouver, Canada Robert Kay, West Vancouver, Canada

**ASSIGNEE:** 

Monsanto Company, St. Louis, MO (U.S. corp.)

APPL-NO:

08/209,752

DATE FILED:

Mar. 9, 1994

ART-UNIT:

184

PRIM-EXMR:

Patricia R. Moody

LEGAL-REP:

Grace L. Bonner, Dennis R. Hoerner, Richard H. Shear

US PAT NO:

5,338,663 [IMAGE AVAILABLE]

L1: 11 of 29

DATE ISSUED: Aug. 16, 1994

TITLE:

Method of identifying inhibitors of .beta.-protein

esterase activity

**INVENTOR:** 

Huntington Potter, Boston, MA

Usamah Kayyali, Somerville, MA

ASSIGNEE:

President and Fellows of Harvard College, Cambridge, MA

(U.S. corp.)

APPL-NO:

07/819,361

DATE FILED: Jan. 13, 1992

ART-UNIT: 185

PRIM-EXMR: Michael G. Wityshyn

ASST-EXMR: Ralph Gitomer

LEGAL-REP: Hamilton, Brook, Smith & Reynolds

US PAT NO: 5,332,672 [IMAGE AVAILABLE] L1: 12 of 29

DATE ISSUED: Jul. 26, 1994

TITLE: Prevention of ES cell differentiation by ciliary

neurotrophic factor

INVENTOR: Joanne Conover, Tarrytown

George D. Yancopoulos, Tarrytown

ASSIGNEE: Regeneron Pharmaceuticals, Inc., Tarrytown, NY (U.S.

corp.)

APPL-NO: 07/865,878 DATE FILED: Apr. 9, 1992

ART-UNIT: 182

PRIM-EXMR: Robert J. Hill, Jr.

ASST-EXMR: Sally P. Teng LEGAL-REP: Gail M. Kempler

US PAT NO: 5,322,938 [IMAGE AVAILABLE] L1: 13 of 29

DATE ISSUED: Jun. 21, 1994

TITLE: DNA sequence for enhancing the efficiency of transcription

INVENTOR: Joan C. McPherson, Vancouver, Canada

Robert Kay, West Vancouver, Canada

ASSIGNEE: Monsanto Company, St. Louis, MO (U.S. corp.)

APPL-NO: 07/977,600 DATE FILED: Nov. 17, 1992

ART-UNIT: 184

PRIM-EXMR: Patricia R. Moody

LEGAL-REP: Grace L. Bonner, Dennis R. Hoerner, Richard H. Shear

US PAT NO: 5,286,487 [IMAGE AVAILABLE] L1: 14 of 29

DATE ISSUED: Feb. 15, 1994

TITLE: Covalent angiogenin/RNase hybrids

INVENTOR: Bert L. Vallee, Brookline, MA

Michael D. Bond, Brighton, MA

ASSIGNEE: President and Fellows of Harvard College, Cambridge, MA

(U.S. corp.)

APPL-NO: 07/953,555

DATE FILED: Sep. 29, 1992

ART-UNIT: 184

PRIM-EXMR: Robert A. Wax ASST-EXMR: Keith D. Hendricks

LEGAL-REP: Allegretti & Witcoff, Ltd.

US PAT NO: 5,283,256 [IMAGE AVAILABLE] L1: 15 of 29

DATE ISSUED: Feb. 1, 1994

TITLE: Cholesterol-lowering agents

INVENTOR: Claude Dufresne, East Brunswick, NJ

Josep Guarro, Tarragona, Spain Leeyuan Huang, Watchung, NJ

Yu L. Kong, Edison, NJ

Russell B. Lingham, Watchung, NJ Maria S. Meinz, Somerset, NJ Keith C. Silverman, Somerset, NJ

Sheo B. Singh, Edison, NJ

ASSIGNEE: Merck & Co., Inc., Rahway, NJ (U.S. corp.)

APPL-NO: 07/918,727 DATE FILED: Jul. 22, 1992

ART-UNIT: 126

PRIM-EXMR: Nicky Chan

LEGAL-REP: Catherine A. Dolan, Melvin Winokur, Paul D. Matukaitis

US PAT NO: 5,270,332 [IMAGE AVAILABLE] L1: 16 of 29

DATE ISSUED: Dec. 14, 1993

TITLE: Cholesteral lowering agents

INVENTOR: Shieh-Shung T. Chen, Morganville, NJ

Leeyuan Huang, Watchung, NJ John G. MacConnell, Westfield, NJ Jon D. Polishook, Scotch Plains, NJ Raymond F. White, Englishtown, NJ

ASSIGNEE: Merck & Co., Inc., Rahway, NJ (U.S. corp.)

APPL-NO: 07/934,134 DATE FILED: Aug. 21, 1992

ART-UNIT: 126

PRIM-EXMR: Nicky Chan

LEGAL-REP: Catherine A. Dolan, Melvin Winokur, Paul D. Matukaitis

US PAT NO: 5,270,204 [IMAGE AVAILABLE] L1: 17 of 29

DATE ISSUED: Dec. 14, 1993

TITLE: Covalent angiogenin/RNase hybrids INVENTOR: Bert L. Vallee, Brookline, MA

Michael D. Bond, Brighton, MA

ASSIGNEE: The President and Fellows of Harvard College, Cambridge,

MA (U.S. corp.)

APPL-NO:

07/947,363

DATE FILED:

Sep. 18, 1992

**ART-UNIT:** 

184

PRIM-EXMR:

Robert A. Wax

**ASST-EXMR:** 

Keith D. Hendricks

LEGAL-REP:

Allegretti & Witcoff, Ltd.

US PAT NO:

5,258,401 [IMAGE AVAILABLE]

L1: 18 of 29

DATE ISSUED: Nov. 2, 1993

TITLE:

Cholesterol lowering compounds

**INVENTOR:** 

Gregory D. Berger, Belle Mead, NJ

Robert W. Marquis, Jr., Iselin, NJ Albert J. Robichaud, Stirling, NJ

Edward M. Scolnick, Wynnewood, PA

**ASSIGNEE:** 

Merck & Co., Inc., Rahway, NJ (U.S. corp.)

APPL-NO:

07/938,981

DATE FILED:

Sep. 10, 1992

**ART-UNIT:** 

126

PRIM-EXMR:

Nicky Chan

LEGAL-REP:

Charles M. Caruso, Melvin Winokur, Carol S. Quagliato

US PAT NO:

5,223,482 [IMAGE AVAILABLE]

L1: 19 of 29

DATE ISSUED: Jun. 29, 1993

TITLE:

Recombinant Alzheimer's protease inhibitory amyloid

protein and method of use

**INVENTOR:** 

James W. Schilling, Jr., Palo Alto, CA

Phyllis A. Ponte, Mountain View, CA

Barbara Cordell, Palo Alto, CA

**ASSIGNEE:** 

Scios Nova Inc., Mountain View, CA (U.S. corp.)

APPL-NO:

07/361,912

DATE FILED:

Jun. 6, 1989

**ART-UNIT:** 

182

PRIM-EXMR:

Robert J. Hill, Jr.

ASST-EXMR:

Nina Ossanna

LEGAL-REP:

Karl Bozicevic

US PAT NO:

5,220,013 [IMAGE AVAILABLE]

L1: 20 of 29

DATE ISSUED: Jun. 15, 1993

TITLE:

DNA sequence useful for the detection of Alzheimer's

disease

**INVENTOR:** 

Phyllis A. Ponte, Mountain View, CA

Barbara Cordell, Palo Alto, CA

**ASSIGNEE:** 

Scios Nova Inc., Mountain View, CA (U.S. corp.)

APPL-NO:

07/444,118

DATE FILED:

Nov. 30, 1989

**ART-UNIT:** 

187

PRIM-EXMR:

Amelia Burgess Yarbrough

LEGAL-REP:

Morrison & Foerster

US PAT NO:

**5,196,525** [IMAGE AVAILABLE]

L1: 21 of 29

DATE ISSUED: Mar. 23, 1993

TITLE:

DNA construct for enhancing the efficiency of

transcription

INVENTOR:

Joan C. McPherson, Vancouver, Canada

Robert Kay, West Vancouver, Canada

ASSIGNEE:

University of British Columbia, Vancouver, Canada (foreign

corp.)

APPL-NO:

07/682,049

DATE FILED:

Apr. 8, 1991

**ART-UNIT**:

184

PRIM-EXMR:

Elizabeth C. Weimar

**ASST-EXMR:** 

P. Rhodes

LEGAL-REP:

Barbara Rae-Venter

US PAT NO:

5,164,316 [IMAGE AVAILABLE]

L1: 22 of 29

L1: 23 of 29

DATE ISSUED: Nov. 17, 1992

TITLE:

DNA construct for enhancing the efficiency of

transcription

**INVENTOR:** 

Joan C. McPherson, Vancouver, Canada

Robert Kay, Vancouver, Canada

**ASSIGNEE:** 

The University of British Columbia, Vancouver, Canada

(foreign corp.)

APPL-NO:

07/395,155

DATE FILED:

Aug. 17, 1989

**ART-UNIT:** 

184

PRIM-EXMR:

Elizabeth C. Weimar

**ASST-EXMR:** 

P. Rhodes

LEGAL-REP:

Barbara Rae-Venter, Bertram I. Rowland

US PAT NO:

5,135,915 [IMAGE AVAILABLE]

DATE ISSUED: Aug. 4, 1992

TITLE:

Method for the treatment of grafts prior to

transplantation using TGF-.beta.

**INVENTOR:** 

Christine W. Czarniecki, San Francisco, CA

Michael A. Palladino, Foster City, CA

Eli Shefter, San Francisco, CA

ASSIGNEE: Genentech, Inc., South San Francisco, CA (U.S. corp.)

APPL-NO: 07/258,276 DATE FILED: Oct. 14, 1988

ART-UNIT: 181

PRIM-EXMR: Merrell C. Cashion, Jr. ASST-EXMR: Andrew G. Rozycki

LEGAL-REP: Janet E. Hasak

US PAT NO: 5,135,849 [IMAGE AVAILABLE] L1: 24 of 29

DATE ISSUED: Aug. 4, 1992

TITLE: In-vitro methods for identifying compositions which are

agonists and antagonists of androgens

INVENTOR: Ana M. Soto, Boston, MA

Carlos Sonnenschein, Boston, MA

ASSIGNEE: Trustees of Tufts College, Medford, MA (U.S. corp.)

APPL-NO: 07/339,800

DATE FILED: Apr. 18, 1989

ART-UNIT: 182

PRIM-EXMR: David A. Saunders LEGAL-REP: David Prashker

US PAT NO: 5,087,368 [IMAGE AVAILABLE] L1: 25 of 29

DATE ISSUED: Feb. 11, 1992

TITLE: Purified protease nexin

INVENTOR: Randy W. Scott, Sunnyvale, CA

Joffre B. Baker, El Granada, CA

ASSIGNEE: Incyte Pharmaceuticals, Palo Alto, CA (U.S. corp.)

University of Kansas, Lawrence, KS (U.S. corp.)

APPL-NO: 07/577,887 DATE FILED: Sep. 5, 1990

ART-UNIT: 136

PRIM-EXMR: Ernest G. Therkorn LEGAL-REP: Morrison & Foerster

US PAT NO: 5,006,252 [IMAGE AVAILABLE] L1: 26 of 29

DATE ISSUED: Apr. 9, 1991

TITLE: Purified protease nexin

INVENTOR: Randy W. Scott, Sunnyvale, CA

Joffre B. Baker, El Granada, CA

ASSIGNEE: Invitron, St. Louis, MO (U.S. corp.)

University of Kansas, Lawrence, KS (U.S. corp.)

APPL-NO: 07/378,434

DATE FILED: Jul. 10, 1989

ART-UNIT: 136

PRIM-EXMR: Ernest G. Therkorn

LEGAL-REP: Irell & Manella

US PAT NO: 4,931,373 [IMAGE AVAILABLE] L1: 27 of 29

DATE ISSUED: Jun. 5, 1990

TITLE: Stable DNA constructs for expression of .alpha.-1

antitrypsin

INVENTOR: Glenn Kawasaki, Seattle, WA

Leslie Bell, Seattle, WA

ASSIGNEE: ZymoGenetics, Inc., Seattle, WA (U.S. corp.)

APPL-NO: 06/663,315 DATE FILED: Oct. 22, 1984

ART-UNIT: 185

PRIM-EXMR: Robin Teskin LEGAL-REP: Seed and Berry

US PAT NO: 4,912,136 [IMAGE AVAILABLE] L1: 28 of 29

DATE ISSUED: Mar. 27, 1990

TITLE: Uses of a substituted 2-phenoxyphenylacetic acid as an

immunosuppressant drug

INVENTOR: Elizabeth M. Wood, Lubnaig, 442 Blackness Road, Dundee,

United Kingdom, DD2 1TQ

APPL-NO: 07/212,915

DATE FILED: Jun. 29, 1988

ART-UNIT: 125

PRIM-EXMR: Stanley J. Friedman

LEGAL-REP: Florence U. Reynolds

US PAT NO: 4,806,471 [IMAGE AVAILABLE] L1: 29 of 29

DATE ISSUED: Feb. 21, 1989

TITLE: Plasmids with conditional uncontrolled replication

behavior

INVENTOR: Soren Molin, Holte, Denmark

Janice A. Light, Henley-on-Thames, United Kingdom

Jens E. L. Larsen, Jordlose, Denmark

ASSIGNEE: A/S Alfred Benzon, Copenhagen, Denmark (foreign corp.)

APPL-NO: 06/610,765

DATE FILED: May 16, 1984

ART-UNIT: 185

PRIM-EXMR: Thomas G. Wiseman

ASST-EXMR: S. Seidman

LEGAL-REP: Bryan, Cave, McPheeters & McRoberts

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ALL L# QUERIES AND ANSWER SETS ARE DELETED AT LOGOFF
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U.S. Patent & Trademark Office LOGOFF AT 12:05:35 ON 06 AUG 95